Field Name	Field Description
Prior Authorization	Blincyto
Group Description	
Drugs	Blincyto (blinatumomab)
Covered Uses	Medically accepted indications are defined using the following sources:
	the Food and Drug Administration (FDA), Micromedex, American
	Hospital Formulary Service (AHFS), United States Pharmacopeia Drug
	Information for the Healthcare Professional (USP DI), the Drug
	Package Insert (PPI), or disease state specific standard of care
	guidelines.
Exclusion Criteria	N/A
Required Medical Information	See "Other Criteria"
Age Restriction	N/A
Prescriber	Prescribed by or in consultation with an oncologist/hematologist
Restrictions	
Coverage Duration	The request will be approved for up to a 12 month duration.
Other Criteria	Initial Authorization:
	Patient has a diagnosis of one of the following forms of Acute
	Lymphoblastic Leukemia (ALL):
	a) Relapsed CD19-positive B-cell precursor ALL
	b) Refractory CD19-positive B-cell precursor ALL
	c) CD19-positive B-cell precursor ALL in first or second
	complete remission with minimal residual disease
	(MRD) greater than or equal to 0.1%
	Provider attests to monitor patient for Cytokine Release
	Syndrome (CRS) and neurological toxicities
	Reauthorization:
	<ul> <li>Patient has a diagnosis of relapsed or refractory CD19-positive</li> </ul>
	B-cell precursor ALL and has not exceeded 9 total cycles of
	Blincyto therapy
	Provider attests to treatment response or stabilization of
	disease
	<ul> <li>Prescriber attests to monitor patient for Cytokine Release</li> </ul>
	Syndrome (CRS) and neurological toxicities
	***For CD19-positive B-cell precursor ALL with MRD, reauthorization
Revision/Review	is not allowed***
Date	Medical Director/clinical reviewer must override criteria when,
4/2024	in his/her professional judgement, the requested item is medically
	necessary.
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